COLLINS et al.

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- 12. (Once Amended) The method of claim 1, wherein said second process is executed using a sequence comparison algorithm [BLAST algorithm].
- 27. (Once Amended) The method of claim 18, wherein the identification of repeat sequences within said genomic region is performed using [Repeat Masker software] a software program that screens sequences for:
- i. interspersed repeats that are known to exist in mammalian genomes and;
  - ii. low complexity DNA sequences.
- 28. (Once Amended) The method of claim 18, wherein the comparison of said at least one repeat sequence-free subsequence with said genome database is performed using a sequence comparison algorithm [BLAST algorithm].
- 40. (New) The method of claim 1, wherein the repeat-free subsequences are each at least 100 bp long.
- 41. (New) The method of claim 18, wherein the repeat-free subsequences are each at least 100 bp long.
- 42. (New) The computer program of claim 34, wherein each nucleotide sequence that does not contain any of the repeat sequences is at least 100 bp long.

## **REMARKS**

## I. Status of the Claims

Claims 11-12 and 27-28 are amended and claims 40-42 are added with this Amendment. Claim 39 is canceled without prejudice to further prosecution. Therefore, claims 1-38 and 40-42 are pending with entry of this Amendment. Appendix A contains a marked-up copy of the amendments. For the convenience of the Examiner, Appendix B provides a list of all claims pending with entry of this Amendment.